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Sarah Ameziane-Le Hir, Céline Raguénès-Nicol, Gilles Paboeuf, Aurélie Nicolas, Elisabeth Le Rumeur, et al.. Cholesterol favors the anchorage of human dystrophin repeats 16 to 21 in membrane at physiological surface pressure.. *Biochimica et Biophysica Acta:Biomembranes*, 2014, 1838 (5), pp.1266-73. 10.1016/j.bbamem.2014.01.010 . hal-00965782

HAL Id: hal-00965782

<https://hal.science/hal-00965782>

Submitted on 13 Jun 2014

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Cholesterol favors the anchorage of human dystrophin repeats 16 to 21 in membrane at physiological surface pressure

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KEYWORDS

Atomic force microscopy, Langmuir film, muscular dystrophy, protein lipid interaction, CRAC sequence, spectrin superfamily

ABSTRACT

Dystrophin (DYS) is a filamentous protein that connects the cytoskeleton and the extracellular matrix via the sarcolemma, conferring resistance to muscular cells. In this study, interactions between the DYS R16-21 fragment and lipids were examined using Langmuir films made of anionic and zwitterionic lipids. The film fluidity was modified by the addition of 15% cholesterol. Whatever the lipid mixture examined, at low surface pressure (20 mN/m) few differences appeared on the protein insertion and the presence of cholesterol did not affect the protein/lipid interactions. At high surface pressure (30 mN/m), the protein insertion was very low and occurred only in zwitterionic films in the liquid-expanded phase. In anionic films, electrostatic interactions prevented the protein insertion outright, and caused accumulation of the protein on the hydrophilic part of the monolayer. Addition of cholesterol to both lipid mixtures drastically modified the protein-lipid interactions: the DYS R16-21 insertion increased and its organization in the monolayer appeared to be more homogeneous. The presence of accessible cholesterol recognition amino-acid consensus sequences in this fragment may enhance the protein/membrane binding at physiological lateral pressure. These results suggest that the anchorage of dystrophin to the membrane *in vivo* may be stabilized by cholesterol-rich nano-domains in the inner leaflet of sarcolemma.

FIGURE 1

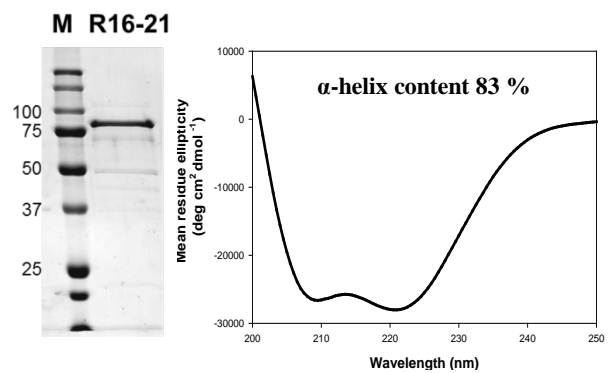


FIGURE 2

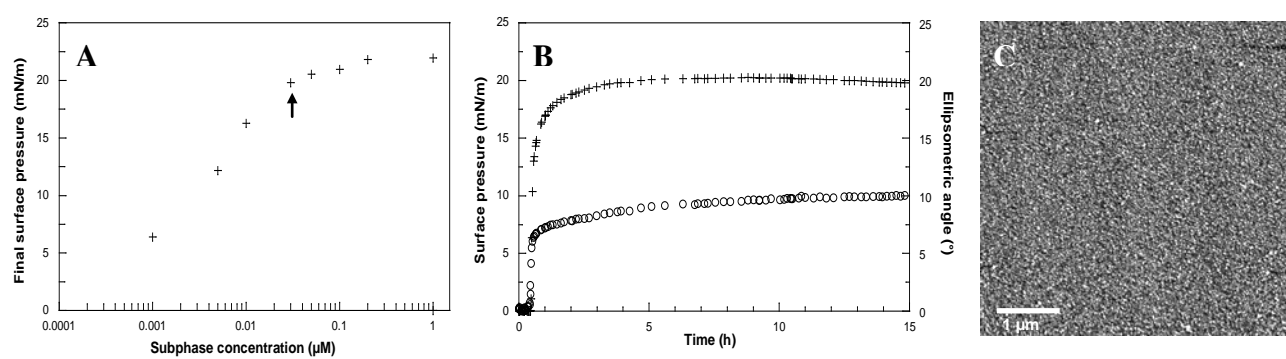


FIGURE 3

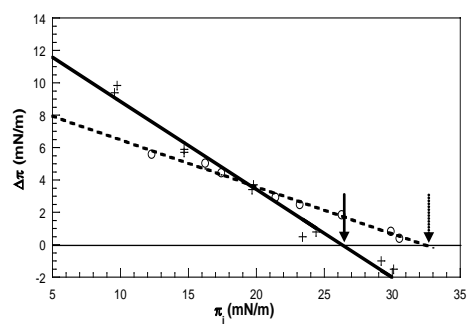


FIGURE 4

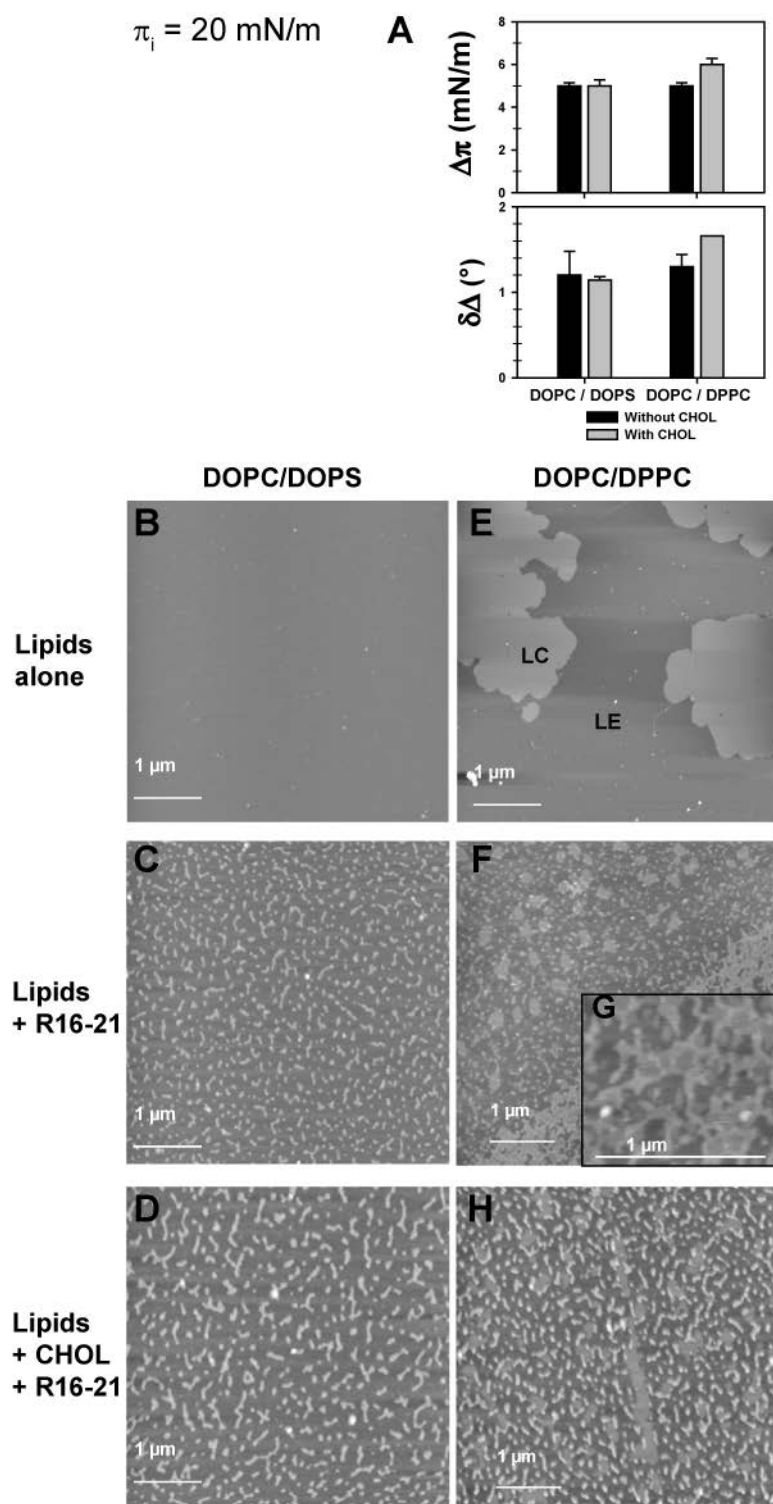


FIGURE 5

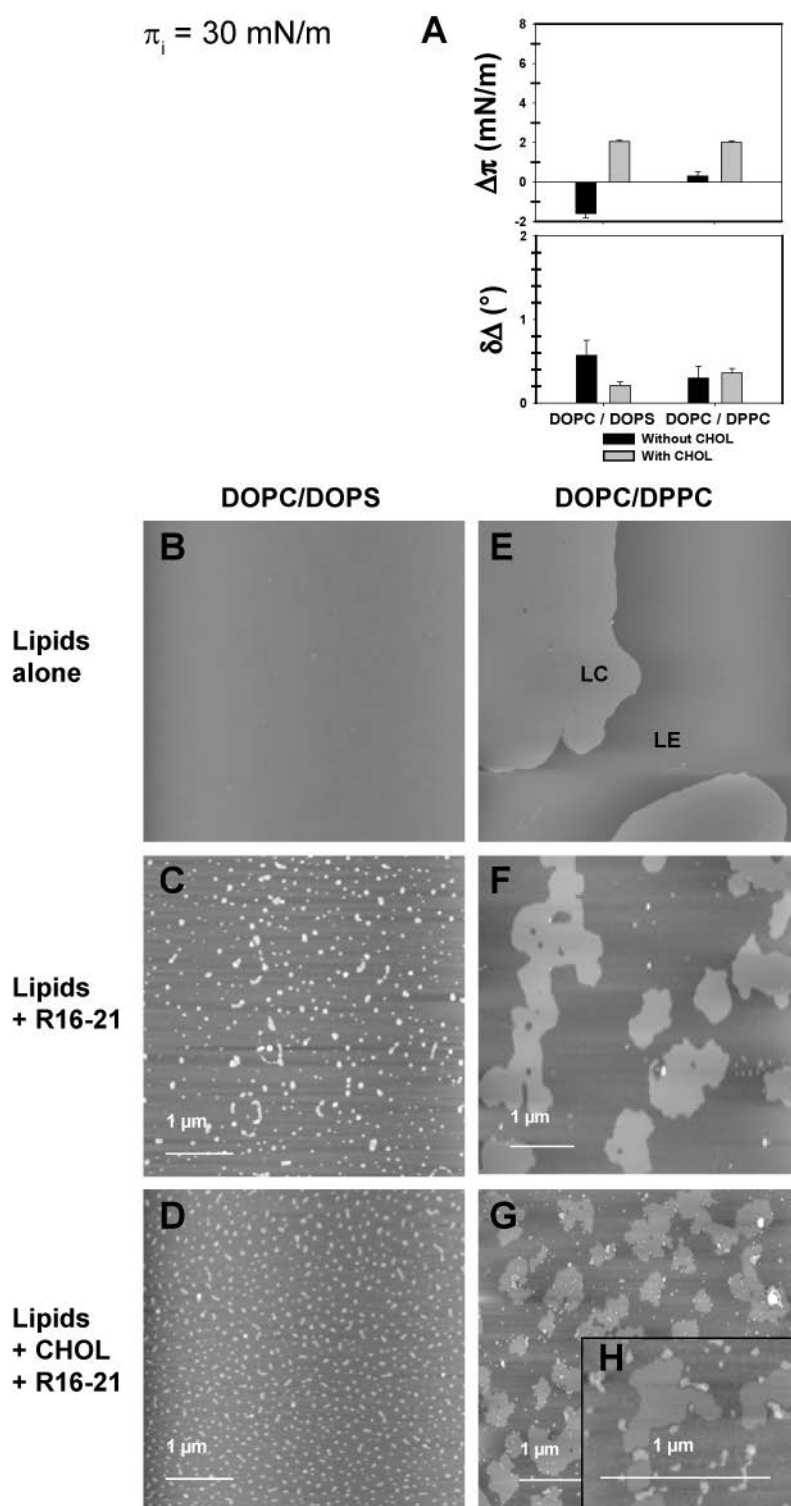
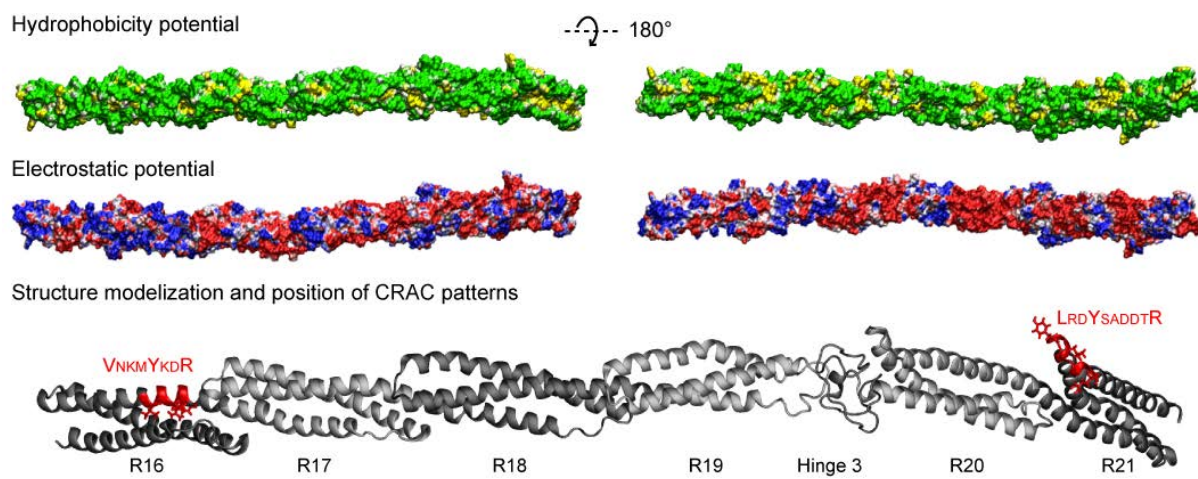


FIGURE 6



SUPPLEMENTARY TABLE S1

Summary of values obtained after protein insertion in lipid monolayers and described in the text: surface pressure ($\Delta\pi$) and ellipsometric angles ($\delta\Delta$) variations; apparent height of protein protrusions in AFM images.

$\pi_i = 20 \text{ mN/m}$

	$\Delta\pi$		$\delta\Delta$		Height (nm)	
	- CHOL	+ CHOL	- CHOL	+ CHOL	- CHOL	+ CHOL
DOPC/DOPS	5 ± 0.1	5 ± 0.3	1.2 ± 0.3	1.1 ± 0.1	2.1 ± 0.2	2.7 ± 0.1
DOPC/DPPC	5 ± 0.1	5.9 ± 0.3	1.3 ± 0.1	1.7	2 ± 0.1	3.1 ± 0.2

$\pi_i = 30 \text{ mN/m}$

	$\Delta\pi$		$\delta\Delta$		Height (nm)	
	- CHOL	+ CHOL	- CHOL	+ CHOL	- CHOL	+ CHOL
DOPC/DOPS	-1.6 ± 0.2	2.0 ± 0.1	0.6 ± 0.2	0.2 ± 0.1	3.6 ± 0.4	2.3 ± 0.1
DOPC/DPPC	0.3 ± 0.2	2.0 ± 0.1	0.3 ± 0.1	0.4 ± 0.1	1.1 à 4	2.6 ± 0.2

SUPPLEMENTARY TABLE S2

Occurrences of CRAC sequences (L/V)–(X)_{1–5}–Y–(X)_{1–5}–(R/K) in the primary sequences of dystrophin (uniprot P11532), alpha erythroid spectrin (uniprot P02549), and beta erythroid spectrin (uniprot P11277) as determined by protein pattern find tool of Sequence Manipulation Suite (Stothard, 2000. Biotechniques 28:1102-1104). Amino acids of CRAC sequences are in blue, sometimes several patterns are overlapped.

DYSTROPHIN		SPECTRIN A	SPECTRIN B
ABD 1	V ₇ EDC Y ER ₁₃ V ₂₁₈ DTTYPDK K ₂₂₆ V ₂₈₀ SLAQQ Y ER ₂₈₈		ABD V ₁₀ GNQPPYSRINAR ₂₂ SH1 - SH2 -
Hinge 1	-	Repeat 1 V ₃₁ LTRYQSFKER ₄₁	Repeat 1 V ₂₇₀ VAFYHYFSKM K ₂₈₁
Repeat 1	-	Repeat 2 L ₄₉ EDSYHLQV F ₅₉	L ₃₄₁ QAFSTYRTVE K ₃₅₂
Repeat 2	-	Repeat 3 -	Repeat 2 L ₃₉₉ EEAEYRR ₄₀₆ L ₄₈₈ EKENYHDQ K ₄₉₈
Repeat 3	-	Repeat 4 V ₂₈₆ LTS E DY G K ₂₉₄ L ₃₅₈ ATSR Y E K ₃₆₅	Repeat 3 -
Hinge 2	-	Repeat 5 L ₃₆₆ QATYW Y HR ₃₇₄	Repeat 4 L ₆₆₃ DY G K ₆₆₇
Repeat 4	-	Repeat 6 L ₅₄₂ IGDDHYDSENI K ₅₅₄	Repeat 5 -
Repeat 5	-	Repeat 7 V ₇₁₀ TSEDY G K ₇₁₇	Repeat 6 -
Repeat 6	L ₉₆₇ SVT D YEIMEQ R ₉₇₈ L ₉₉₆ YYLSTTV K ₁₀₀₄	Repeat 8 -	Repeat 7 -
Repeat 7	-	Repeat 9 -	Repeat 8 -
Repeat 8	L ₁₂₄₃ TTN Y QWLCT R ₁₂₅₃	Repeat 10 -	Repeat 9 V ₁₂₃₅ AEGNLYSDKI K ₁₂₄₆
Repeat 9	L ₁₂₇₁ LS Y LEKANK ₁₂₈₀	SH3 V ₉₈₂ MAL Y DFQAR ₉₉₁ V ₁₀₂₆ PAVYVR R ₁₀₃₃	Repeat 10 V ₁₃₀₀ SYDEAR ₁₃₀₆
Repeat 10	L ₁₃₉₃ AA I AD K ₁₄₀₀	Repeat 11 -	Repeat 11 -
Repeat 11	L ₁₅₄₈ KLHYNELG A K ₁₅₅₈	Repeat 12 -	Repeat 12 -
Repeat 12	L ₁₆₇₁ NLL L EY Q K ₁₆₇₉	Repeat 13 -	Repeat 13 V ₁₆₃₃ EDYGRNI K ₁₆₄₁
Repeat 13	-	Repeat 14 L ₁₄₆₀ IADEH Y AK ₁₄₆₈	Repeat 14 V ₁₆₆₈ DKHYAGL K ₁₆₇₆ L ₁₆₈₆ ENMYHLFQL K ₁₆₉₆
Repeat 14	-	Repeat 15 L ₁₄₉₆ GDYANL K ₁₅₀₃	Repeat 15 L ₁₇₉₂ LAAS Y DLHR ₁₈₀₁
Repeat 15	-	Repeat 16 -	Repeat 16 L ₁₈₆₂ QTAYAGE K ₁₈₇₀
Repeat 16	V ₂₀₈₈ NKM Y KDRQGR ₂₀₉₈	Repeat 17 V ₁₇₃₆ SSQDY G R ₁₇₄₃	Repeat 17 -
Repeat 17	-	Repeat 18 -	
Repeat 18	-	Repeat 19 -	
Repeat 19	-	Repeat 20 -	
Hinge 3	-	Repeat 21 -	
Repeat 20	-	EF-hand 1 -	
Repeat 21	L ₂₆₄₅ KLLRD Y SADD T R ₂₆₅₇	EF-hand 2 -	
Repeat 22	-	EF-hand 3 L ₂₃₆₄ AEGKS Y IT K ₂₃₇₃	
Repeat 23	L ₂₈₈₁ EKL Y QEPR ₂₈₈₉		
Repeat 24	-		
Hinge 4	V ₃₁₀₁ RFSA Y RTAM K ₃₁₁₁		
Cys-rich	L ₃₁₅₄ TTI Y DR ₃₁₆₀ L ₃₁₈₂ LN V YDTGR ₃₁₉₀ L ₃₂₁₁ EDKYR Y LF K ₃₂₂₀ L ₃₅₂₀ QAE Y DRL K ₃₅₂₈		
C-term			

1. Introduction

Dystrophin (DYS) is a long filamentous muscular protein of 427 kDa (about 175 nm long) [1, 2]. Most of this length is due to a long central rod domain composed of 24 spectrin-like repeats (R1 to R24) folded into a triple- α -helical bundle [3]. The biological function of dystrophin is to protect the membrane against stress during elongation/contraction muscular cycles. Its absence leads to the severe disease Duchenne muscular dystrophy (DMD), whereas the milder Becker muscular dystrophy (BMD) may be observed when truncated proteins are expressed [4]. Due to its high levels of STRs, dystrophin is a member of the spectrin-like superfamily [5]. A relationship has been suggested between the structural organization of the repeat domains in these proteins and the membrane's protection against shearing stresses [3]. This function may result from a homogeneous protein distribution along the membrane caused by protein/lipid interactions. Indeed, spectrin is known to bind to the inner membrane leaflet due to interactions with phosphatidylserine [6-9]. Moreover the presence of 10-20% cholesterol (CHOL) facilitates the insertion of spectrin into phosphatidylethanolamine/phosphatidylcholine monolayers that were preformed at low surface pressure, i.e. from 8 to 11 mN/m [10]. Even during its liquid-expanded phase (LE), spectrin binding capacity depends on the fluidity of the monolayer. Our previous studies showed that fragments of the dystrophin rod domain interact with membrane system models through electrostatic and/or hydrophobic forces [11, 12].

This study is focused on the interaction of the specific dystrophin fragment made of repeats R16 to R21 and a membrane model. This construct has several specific features that set it apart from other dystrophin fragments. For instance, it contains a hinge (H3) between the repeats R19 and R20 [13, 14]. The presence of H3 in microdystrophin improves the capacity for muscle degeneration prevention, while deletion of this hinge leads to the BMD phenotype [15, 16]. In addition, *DYS* R16-21 is coded by exons 42 to 53, which includes the hot-spot between exons 45 to 53 where about one third of BMD patients have mutations [17]. The fragment also mediates specific protein interactions, such as the anchoring of neuronal NO Synthase (nNOS) to the sarcolemma through R16-R17 [18]. The mislocalisation of nNOS in the membrane leads to a decrease of muscular strength in dystrophin-deficient *mdx* mice [19]. R16-21 is located in the last part of the protein, where the cysteine-rich domain is known to bind to β -dystroglycan (β -DG) in a membrane protein complex [20] that is located in detergent-resistant membrane. Sarcolemmal CHOL depletion induces the weakening of force contractions through the redistribution of β -DG, and is accompanied by a reduction of

β -DG/dystrophin interactions in favor of β -DG/caveolin-3 interactions [21]. All these studies showed that structural or lipid environmental modifications impact muscular activity. Nevertheless, up to now little was known about the ability of DYS R16-21 to interact with lipid membranes.

Here, we address the question of the DYS R16-21 anchorage in mixed lipid films and more particularly of the effect of CHOL on the interaction. Since phosphatidylserine is specifically abundant in the inner leaflet of the sarcolemma membrane, the effect of the polar head charge was tested using a simplified mixture composed of anionic dioleoylphosphatidylserine (DOPS) and zwitterionic dioleoylphosphatidylcholine (DOPC) lipids. Because we chose unsaturated lipids, the monolayers are in the LE phase regardless of the surface pressure. A mix of dipalmitoylphosphatidylcholine (DPPC) and DOPC was used to mimic a detergent-resistant membrane as described [22, 23]. Indeed DPPC has been reported to form liquid-condensed (LC) phase domains and also to create separated phases when mixed with DOPC [24]. Cholesterol, also present in the inner leaflet up to 25% [25], is known to modulate membrane fluidity, and particularly at high surface pressure, it acts as a head group spacer because of its molecular orientation in the monolayer. Moreover, CHOL is located mainly in the condensed phase formed by DPPC [26, 27]. Langmuir monolayers were used, which allowed for control of charge densities, of fluidity, and of lipid packing according to specific lipid compositions and surface pressures. Protein-lipid interactions and protein structure were determined by combining tensiometric and ellipsometric measurements with atomic force microscopy imaging. We evaluated the influence of 15% CHOL on the interaction of DYS R16-21 with DOPC/DOPS and DOPC/DPPC lipid mixtures. Using two initial surface pressures, two different lipid packings were tested. The higher of these (30 mN/m) is considered to be comparable to physiological membrane pressure [28, 29].

Our results show that DYS R16-21 interacts with the lipid monolayer no matter what mixture was used, while the surface pressure has a notable impact on the insertion. In addition, under high surface pressure CHOL was shown to stabilize the fragment at lipid interface, likely through the presence of two cholesterol recognition amino-acid consensus (CRAC) sequences on the protein surface [30]. The presence of CRAC sequences all along dystrophin allows specific protein/lipid interactions that result in homogeneous protein distribution along the membrane. This anchorage could be related to lateral force transmission during muscle elongation/contraction.

2 Experimental sections

2.1 Materials

All lipids were purchased from Avanti Polar Lipids (Alabama, US). The pGEX-4T-1 plasmid vector and GSTrap HP and Sephacryl S-100 columns were bought from GE Healthcare. BL21 bacteria and restriction enzymes were purchased from Ozyme (St.-Quentin-en-Yvelines, France). Purified bovine thrombin came from Stago (Asnières-sur-Seine, France).

2.2 Preparation of recombinant DYS R16-21 protein

The DYS R16-21 fragment begins at residue Ser 1994 and ends with Thr 2686, using the full-length human dystrophin annotation (NCBI NP_003997). In order to improve stability and as previously done for other dystrophin fragments [31, 32], we included a three-amino-acid extension in the N-terminus and a seven-amino-acid extension in the C-terminus. The protein was cloned in pGEX-4T-1 then expressed in a protease-deficient *E. coli* BL21 strain as a GST-tagged protein [33]. Cell lysis was obtained by sonication after lysozyme treatment. DYS R16-21 was further purified using GST affinity chromatography. After thrombin proteolysis (50 UI for 2 hours at room temperature), the chromatography fractions were separated by salting-out with ammonium sulfate. The target protein was collected at 24% salt saturation and rediluted in TNE buffer (20 mM Tris, 150 mM NaCl, 0.1 mM EDTA, pH 7.5). Purification was achieved with size-exclusion chromatography by passing it through a Sephacryl S-100 column (L 2.6 cm x H 60 cm) at 1 mL/min using the same buffer. After purification, the protein was concentrated using a 30 kDa cut-off centrifugal concentrator. Purity was assessed by 10% SDS-Polyacrylamide gel electrophoresis (SDS-PAGE), and concentration was determined using a bicinchoninic acid assay with a standard of serum albumin.

Circular dichroism spectra were acquired between 250 nm and 200 nm, with 1.5 μ M of protein in TNE buffer at 20°C (J-815, Jobin-Yvon) using 0.2 cm path-length cells. The α -helical content was calculated from the mean molar residue ellipticity $[\theta]$ (mdeg.cm².dmol⁻¹) at 222 nm [34].

2.3 Tensiometric and ellipsometric measurements

In order to investigate the interfacial properties of the protein with and without lipids, surface pressure and ellipsometric angles were measured in Teflon troughs (4, 8, or 60 mL) at 20°C. The Wilhelmy method was used to determine the surface pressure using a tensiometer (Nima Technology, Cambridge, UK). Ellipsometric measurements were performed with a conventional polarizer-analyzer null-ellipsometer setup. Briefly, a He-Ne laser (632.8 nm;

Melles Griot, Carlsbad, CA) with an angle of incidence of 52.18° (1° away from the Brewster's angle), was used as the light source. Reflecting surface properties were explored by recording the positions of the polarizer and analyzer having the minimal intensity of transmitted light. The analyzer angle was doubled to yield the value of the ellipsometric angle (Δ) [35-37]. Both Δ and the surface pressure (π) were recorded as functions of time. The amphiphilic character was determined by reporting the surface pressure reached at this end of the absorption kinetics as a function of the protein's subphase concentrations (ranging from $0.0001 \mu\text{M}$ to $1 \mu\text{M}$). For lipid/protein experiments, the lipid mixtures (DOPC/DOPS 1:1 molar ratio and DOPC/DPPC 1:1 molar ratio) were prepared in a 2:1 chloroform/methanol solution (v/v) at 0.5 mM , and where used, CHOL was added at 15% (molar ratio). The lipids were gently spread at the air/liquid interface of the TNE subphase at the desired surface pressure. These initial lipid surface pressures (π_i) ranged from 5 to 35 mN/m. At a final concentration of $0.03 \mu\text{M}$, the protein was then injected into the subphase just beneath the lipid monolayer. The variations ($\delta\Delta$ and $\Delta\pi$) induced by protein adsorption at the end of the absorption kinetics were reported as functions of time and π_i . Each experiment was repeated up to four times and means \pm standard deviation (SD) were provided for each series. The temperature varied between 19 and 21°C .

2.4 Atomic force microscopy (AFM) observations

The interfacial film was transferred onto a freshly-cleaved mica plate using the Langmuir-Blodgett technique. The experiment was realized using a 102 cm^2 Langmuir trough equipped with two movable barriers and controlled by a computer (model 601 M, Nima Technology, Cambridge, UK). The transfer was performed at the end of the kinetics and at a constant surface pressure. The dipper speed rate was 1 mm/min . Transferred monolayers were imaged using a PicoPlus atomic force microscope (Agilent Technologies, Phoenix, AZ) equipped with a $10 \mu\text{m}$ scanner operating in contact mode. Images were acquired under ambient conditions using silicon nitride tips on integral cantilevers (ScienTec) with a spring constant of 0.06 N/m . For each measurement, the set point was adjusted before and during the scanning to minimize the force between the tip and the sample. In each case, images were obtained from at least two samples prepared on different days, with at least four macroscopically-separated areas in each sample. To determine the height of the objects, after processing the images with Gwyddion 2.26 software (<http://gwyddion.net>) we did a statistical analysis. Objects of interest were selected by threshold, and then areas and maximal heights were determined. The values of height versus area were plotted, resulting in a plateau. This

plateau included the majority of the objects and was used to compare the different experimental conditions.

2.5 Three-dimensional molecular homology modeling and surface properties

Three-dimensional (3D) homology models were obtained using I-TASSER server [38]. As previously described [39], the sequences of two successive repeats (“tandem repeats”) with an overlap of a single repeat were submitted to I-TASSER (e.g. DYS R16-17, then DYS R17-18, and so on). The H3 sequence was submitted separately. Superposition of the shared repeat of two successive tandem repeats allowed the construction of an extended model which we further minimized by YASARA (e.g. the DYS R16-17 and DYS R17-18 models were used to construct a DYS R16-18 model). In this way, step by step, the global 3D structure was reconstructed. Surface hydrophobicity was calculated using PLATINUM [40] and surface electrostatic potentials in the presence of 50 mM NaCl were calculated using the Adaptive Poisson-Boltzmann Solver (APBS) software [41]. Visualization was done using VMD and PyMOL.

3. Results

3.1 Protein purification

SDS-PAGE of the recombinant human dystrophin fragment DYS R16-21 (figure 1A) showed the protein at the expected molecular weight of 80 kDa. As expected with spectrin-like repeats [11, 42], the circular dichroism spectrum of DYS R16-21 displayed the typical predominant α -helix folding with two minima at 208 and 222 nm (figure 1B). Per Chen et al., the molar ellipticity of 222 nm allowed for determination of an α -helix content of 83% [43]. The ratio of the molar ellipticity at 222 and 208 nm ($\theta_{222}/\theta_{208}$) was 1.05, indicating that the helices were interacting and folded in a coiled-coil [44]. Therefore, it appears that the recombinant DYS R16-21 fragment was highly purified and correctly folded, conditions necessary for its further study.

3.2 Amphiphilic properties of DYS R16-21

For each DYS R16-21 subphase concentration between 0.0001 to 1 μ M, surface pressure measurements were done and the final surface pressure versus subphase protein concentrations are reported in figure 2A. As the protein concentration increased, π increased

until a plateau (π_{\max}) of 21 mN/m was reached, indicating the amphiphilic character of DYS R16-21. From these adsorption kinetics results, the critical concentration required for interface saturation was determined to be 0.1 μM . For further experiments, a protein concentration of 0.03 μM (or 2.5 $\mu\text{g/mL}$) was chosen (indicated with an arrow in figure 2A) to avoid a potential aggregation of proteins in the subphase or the formation of multilayers at the interface. Figure 2B shows a typical adsorption time for this protein concentration. Whereas surface pressure is related to the protein's ability to cover the interface and to interact laterally, the ellipsometric angle depends on the amount of matter at the interface. As is usual, at the very beginning of the adsorption the ellipsometric angle increased quicker than the surface pressure. It reached an angle of 10° at equilibrium, a value corresponding to a monolayer as experienced in other proteins [45]. AFM imagery of this monolayer at the end of the adsorption kinetics (figure 2C) showed a homogeneous film with no aggregation.

3.3 DYS R16-21 interaction with lipid monolayers

The adsorption kinetics of DYS R16-21 was monitored at various initial lipid surface pressures (π_i). The variations of the surface pressure ($\Delta\pi$) i.e. the difference between the initial lipid surface pressure and the final surface pressure were reported as a function of π_i (figure 3). When π_i increased, $\Delta\pi$ induced by adsorption of DYS R16-21 decreased. Indeed as the lipids compacted, the protein was less and less able to be inserted into the lipid monolayer. When there was no more variation between the initial and final surface pressures ($\Delta\pi=0$ or intercept of the linear regression with the x axis as indicated with arrows in figure 3), the maximal insertion pressure (MIP) was reached [46]. This was clearly dependent on the lipid mixture, with values of 26.5 mN/m for DOPC/DOPS and 33 mN/m for DOPC/DPPC.

For π_i less than 20 mN/m, $\Delta\pi$ was higher with anionic than with zwitterionic lipids, suggesting an attractive role for electrostatic interactions in the insertion of DYS R16-21. On the other hand, when π_i was higher than 20 mN/m, $\Delta\pi$ was higher with zwitterionic lipids, suggesting presence of hydrophobic interactions during the protein's insertion process, while electrostatic interactions limit the insertion. Moreover, for the DOPC/DOPS monolayer at an initial surface pressure higher than π_e (26 mN/m), the protein induced a surface pressure decrease ($\Delta\pi = -1.6$ mN/m), thus showing that attractive interactions were still taking place between DYS R16-21 and the compacted DOPC/DOPS monolayer.

To further analyze the organization of DYS R16-21 in lipid environments, we focused our work on lipid monolayers formed at two significant initial surface pressures: 20 mN/m, where

the protein adsorption induced similar overpressures for all lipid compositions tested; and 30 mN/m, considered to be close to physiological membrane pressure [29]. All reported values are recapitulated in table S1.

3.4 Cholesterol has few effects at a π_i of 20 mN/m

Cholesterol is known to influence lipid packing therefore we studied the impact of 15 % CHOL on the DYS R16-21/lipid interactions. At π_i of 20 mN/m (figure 4A), no changes were observed upon the addition of CHOL to the DOPC/DOPS film, while there was a slight increase of $\Delta\pi$ and $\delta\Delta$ (1 mN/m and 0.4° respectively) in the DOPC/DPPC mixture. AFM images of pure lipid films were unmodified in the presence of CHOL. DOPC/DOPS \pm CHOL (figure 4B) formed a homogeneous liquid-expanded (LE) phase. DOPC/DPPC \pm CHOL images (figure 4E) showed the coexistence of an LE phase of DOPC and a liquid-condensed (LC) phase of DPPC, as previously observed [47]. The LC domains were several μm in diameter and their apparent height was 1.4 ± 0.1 nm.

Mixed protein/lipid films showed bright protrusions attributed to the protein's presence (figure 4C-H). In DOPC/DOPS \pm CHOL films, these protrusions appeared mostly as worm-like curved filaments (figure 4C-D). CHOL induced both protein clustering and an increase in the apparent height of protrusions (2.1 ± 0.1 nm without CHOL versus 2.7 ± 0.1 nm with CHOL). These changes in DYS R16-21 organization occurred even though the lateral pressure and the amount of material at the interface were similar with or without CHOL (figure 4A).

In DOPC/DPPC films (figure 4E-H), the presence of DYS R16-21 drastically decreased the LC domain's diameter, which never exceeded 400 nm. Without CHOL (figure 4F-G), the proteins appeared as round objects or filaments with heights higher (2.0 ± 0.1 nm) than those of LC domains (1.4 ± 0.1 nm). The filaments were most often stuck to an LC phase (detailed view, figure 4G) whereas round objects were located in an LE phase. In the presence of CHOL (figure 4H), higher filaments (3.1 ± 0.2 nm) were located both in the LE phase and at the boundaries of LE/LC phases.

3.5 Cholesterol increases DYS R16-21 insertion at a π_i of 30 mN/m

Ellipsometric and tensiometric measurements (figure 5A) showed that in any lipid condition, a π_i of 30 mN/m induced the presence of less protein at the interface than with a π_i of 20 mN/m. Injection of DYS R16-21 under the DOPC/DOPS/CHOL monolayer led to an

increase in surface pressure (from -1.6 mN/m without CHOL to +2 mN/m with CHOL) and a decrease in the ellipsometric angle (from 0.6° without CHOL to 0.2° with CHOL), indicating modified protein/lipid interactions that favor the insertion of DYS R16-21. In the DOPC/DPPC films, the presence of CHOL had no significant effects on ellipsometric angle variation after injection of DYS R16-21, whereas the presence of CHOL induced an increase in $\Delta\pi$ (from 0.3 mN/m without CHOL to 2 mN/m).

The AFM images of DOPC/DOPS films with or without CHOL (figure 5B) were similar to those at 20 mN/m. The DOPC/DPPC/CHOL AFM images revealed that the presence of CHOL caused a slight decrease in height of LC domains (from 1.4 nm without CHOL to 1.2 nm) (figure 5E).

In the DOPC/DOPS/protein films (figure 5C), DYS R16-21 appeared as protrusions forming heterogeneous clusters higher (3.6 ± 0.4 nm) and larger than those previously observed at a π_i of 20 mN/m. This is consistent with the variations in surface pressures and ellipsometric angles seen with the two surface pressures (figure 4A and 5A). In the presence of CHOL (figure 5D), the proteins formed small protrusions that are smaller in length, width, and height ($2.3 \text{ nm} \pm 0.1$) than those in the absence of CHOL or with a smaller π_i . Finally, the protein organization was more homogeneous, with deeper insertion into the lipid monolayer and less material at the interface.

In DOPC/DPPC/protein films (figure 5F), the protein was present as scarce clusters in agreement with the low $\Delta\pi$. Their heights (between 1 and 4 nm), were very heterogeneous. By contrast, CHOL induced numerous clusters of DYS R16-21 (figure 5G-H) with heights of 2.6 ± 0.2 nm. The LC domain borders were much more irregular and angular than without CHOL. The protein clusters appear to be more elongated when in contact with LC domains than when embedded in LE domains (detailed view, figure 5H). Some objects also appeared to be embedded in LC domains, but since there was a depression surrounding these clusters it was obvious that there was no direct contact between the LC and the protein.

3.6 Computational analysis of DYS R16-21 in presence of 50 mM NaCl

As expected for a soluble protein, the molecular surface of DYS R16-21 is mainly hydrophilic. Nevertheless one side of the fragment displays a higher presence of hydrophobic amino acids (figure 6). The electrostatic potential at the molecular surface shows a distribution of negative and positive charges along the molecule. Numerous positive patches were observed, mainly on repeats 16 and 17 of DYS R16-21, with the rest dispersed more

widely along the molecule and embedded in the negative surface. A rotated view of the fragment showed a more contrasted surface with larger negative patches, separated by a few small positive zones that appear mostly on repeats 18 and 19. Interestingly, some hydrophobic patches colocalize or are very close to the cationic surface (for example on R16, R18 and R20). These may thus be anchorage points for interaction or insertion in anionic and zwitterionic lipid monolayers even under high initial surface pressures.

A search of the primary sequence of dystrophin revealed several occurrences of the CRAC pattern $-(L/V)-(X)_{1-5}-Y-(X)_{1-5}-(R/K)-$ [30]. Due to this sequence, a cholesterol binding cleft is created where Y and R/K interact directly with CHOL [48]. In DYS R16-21, the pattern was found in the C helix of R16 ($V_{2088}NKMYKDR_{2098}$) and in the R21 B-C loop ($L_{2647}RDYSADDTR_{2657}$) (figure 6). Most of the amino acids involved are exposed on the surface, and accessible enough to enable interaction with CHOL molecules.

4. Discussion

In this study, we examine the interaction of DYS R16-21 with lipid monolayers. This particular dystrophin fragment is characterized by the presence of hinge 3 (H3) and is frequently mutated in patients. DYS R16-21 displayed amphiphilic properties that were probably due to the presence of hydrophobic amino-acids all along the surface (figure 6). Compared with previously-studied rod domain fragments, the adsorption of DYS R16-21 at the air/water interface was faster, the plateau was reached at a smaller protein concentration but has a similar value of 21 mN/m [12, 49]. This indicates both that the interface is more rapidly covered by the protein, and that the molecules do indeed interact with each other. This result could be related to the larger size of the protein as compared to other dystrophin fragments previously studied, and/or to the presence of H3, which may make the protein's structure more flexible [13].

The monolayer's initial lipid surface pressure (π_i) is a major factor influencing the adsorption of amphiphilic proteins through electrostatic or hydrophobic forces because it determines lipid packing. When π_i was below 20 mN/m, the protein was able to insert itself into both anionic and zwitterionic monolayers; however the maximal insertion pressure was significantly lower for the anionic mixture suggesting that attractive electrostatic interaction plays a major role in DYS R16-21-membrane interactions. The negative charge of the polar head attracts the protein toward the membrane, allowing further hydrophobic interactions to occur thanks to free interfacial space. At a π_i of 20 mN/m, the protein insertion overpressures

and ellipsometric angle variations seem to be equivalent for both lipid mixes. AFM observations show the protein as round or fibrous objects with regular heights (about one protein thick). The fibrous objects were present in the anionic mix, and interconnected filaments were only detected in contact with LC domains. These features imply that the molecules were stabilized at the interface by negative charges or LC domains and then adopted the same orientation at the lipid interface. This favors the existence of lateral protein/protein interactions in both lipid mixtures. At a π_i over 20 mN/m, electrostatic interactions still occur but the space between lipid head groups becomes more and more limited, and the fragment insertion slows. Finally, when the π_i of the DOPC/DOPS film was over 26 mN/m, interactions with lipid head groups caused a reorientation and relaxation of the films, as shown by a decrease in the final surface pressure, and the protein appears as irregular clusters. This phenomenon has been reported previously for the dystrophin R2 repeat [50], DYS R20-24 [12] and in α -spectrin [6]. In zwitterionic film, however, the insertion of DYS R16-21 occurs up to 31 mN/m. This is probably because the protein's orientation is not driven by electrostatic interactions and because its flexibility allows for penetration of the films even at higher surface pressures.

When comparing the anionic monolayer insertion of the previously-studied fragments DYS R1-3, DYS R11-15 and DYS R20-24 to that of the DYS R16-21 studied here, different behaviors are seen, thus demonstrating that these fragments are not interchangeable as had been stated [51]. Indeed, at 30 mN/m, only DYS R11-15 inserts into the monolayer ($\Delta\pi = +2$ mN/m) [49], whereas both DYS R16-21 and R20-24 stayed at the head group level [12]. At 20 mN/m, a gradient seems to occur along the central domain, with greater insertion for fragments from the N- to C-terminal ends ($\Delta\pi = 7$ mN/m for DYS R1-3, 6 mN/m for DYS R11-15, and 5 mN/m for DYS R16-21 and DYS R20-24). This intriguing observation must be further studied to decipher whether it has a physiological significance. It would be especially interesting to understand the relationship between the variation in sarcolemmal lateral pressure during muscle elongation/contraction, and the number or position of anchoring points along the protein. Interestingly, it has been reported that binding of spectrin to phosphatidylserine in the inner membrane increases the mechanical stability of red blood cell membranes [52].

We examined dystrophin/lipid/CHOL interactions because of cholesterol presence in the membrane and of its impact on the membrane's physical properties. Moreover we report the existence of a specific amino-acid sequence called CRAC (cholesterol recognition amino-

acid consensus) and involved in CHOL binding [30, 53]. Like spectrin [54, 55], dystrophin has been shown to be associated *in vivo* to detergent-resistant membranes [56] and the interaction between β -DG and dystrophin is weakened by CHOL depletion [21]. The impact of CHOL on the structural and lateral organization of lipid layers is highly-documented [27, 57, 58], in particular its role as head group spacer and its usual localization in the condensed phase formed by DPPC [26]. In addition, several monolayer experiments have shown complex formation between CHOL and phosphatidylserine, an important component of the inner leaflet [59]. Our study was performed at 15 % of CHOL since 10 to 20 % CHOL have been shown to increase spectrin insertion in phosphatidylethanolamine/phosphatidylcholine monolayers [10] and also because this concentration is known to preserve lateral phase separation [60]. At the lower initial surface pressure (20 mN/m), macroscopic measurements (π , Δ) were unmodified in presence of CHOL, but topographic images revealed an increase in the lateral size and height of protein protrusions, indicating changes in the molecular organization of DYS R16-21 within the lipid monolayers. CHOL favors lateral protein-protein interactions, and since the lipids are in fluid phase, it increases steric packing in the hydrophobic part, limiting deep anchoring. In the DOPC/DPPC mixture, LC domain fragmentation is induced by the protein. DYS R16-21 insertion might induce local disorders in the lateral packing of the acyl chains. These hydrophobic mismatch defects result in lateral fragmentation of these domains, and this phenomenon is amplified by the presence of CHOL [61, 62]. At a higher initial surface pressure ($\pi_i=30$ mN/m), CHOL amplifies protein insertion into lipid monolayers and favors an homogeneous organization of the protein in both LC and LE domains. Indeed, the sterol polar head group is small compared to choline or serine, while the hydrophobic part at this pressure is similar to that of a phosphatidylcholine-saturated chain [24], thus CHOL plays the role of head group spacer. This promotes a deeper and more homogenous lipid insertion. The DOPS/DYS R16-21 electrostatic interactions are modified by CHOL. This could be linked either to PS/CHOL complex formation as suggested by Radhakrishnan *et al* [59] or to specific DYS R16-21/CHOL interactions.

Bio-informatics analysis of the DYS R16-21 amino acid sequence revealed two occurrences of the CRAC sequence $-(L/V)-(X)_{1-5}-Y-(X)_{1-5}-(R/K)-$ [30]. CRAC sequences are also present in spectrin and in other dystrophin fragments (Table S2). These patterns in DYS R16-21 are easily accessible for interaction at the protein surface. Not only has CRAC been identified in many proteins that interact with cholesterol, but it has been more widely suggested to be related to protein propensity for interaction with cholesterol-rich lipid

domains [63]. CRAC motifs are often reported in membrane proteins, but are compatible with peripheral proteins and was also reported in a secreted toxin [64].

5. Conclusion

Our results highlight the fact that dystrophin fragments, and here especially DYS R16-21, can interact directly with monolayers at high surface pressure, at low fluidity, and in the presence of anionic lipids. We further show that the presence of dystrophin close to the membrane is probably not due only to a link with β -DG but also thanks to its own sub-part membrane-binding properties. It has now to be tested whether our *in vitro* results may be transposed *in vivo*. Does formation of direct interactions between dystrophin and cholesterol-rich nano-domains stabilize the dystrophin/ β -DG/sarcolemma in muscular cells? If so, cholesterol-rich domains in the inner sarcolemma layer could prevent ejection of dystrophin under surface pressure variations due to contraction/elongation cycles and contribute to sarcolemma resistance.

FIGURES LEGENDS

Figure 1: DYS R16-21 sub-domain purification and secondary structure analysis. (left) 10 % SDS polyacrylamide gel of DYS R16-21 obtained after purification. Lane M is the protein ladder. (right) Circular dichroism measurements of 1.5 μM of DYS R16-21 in TNE buffer (pH 7.5) at 20°C.

Figure 2: DYS R16-21 adsorption at the air-liquid interface. (A) Final surface pressure reached at the end of the adsorption kinetics versus DYS R16-21 subphase concentrations. Arrow indicates π reached for protein subphase concentration of 0.03 μM (B) Adsorption kinetics of DYS R16-21 at air-liquid interface for the subphase concentration of 0.03 μM , surface pressure (+), and ellipsometric angle (o). (C) AFM topographic image of transferred DYS R16-21 monolayer at the end of the kinetics (5 h). Surface pressure was 20 mN/m. Scan size 5 μm x 5 μm , z scale 10 Å.

Figure 3: Overpressure induced by adsorption of DYS R16-21 at air/lipid interface versus the initial lipid surface pressures of DOPC/DOPS (full line) and DOPC/DPPC (dotted line). Arrows indicate the values of maximal insertion pressures (MIP). The subphase concentration of DYS R16-21 was 0.03 μM .

Figure 4: Characterization of DYS R16-21 at the initial lipid surface pressure of 20 mN/m. (A) Variations of the surface pressure ($\Delta\pi$) and of the ellipsometric angle ($\delta\Delta$) induced by DYS R16-21 adsorption on phospholipid monolayers (black) and phospholipid/CHOL monolayers (gray). (B-H) Typical AFM topographic images of monolayers as indicated on the panels. Scan size 5 μm x 5 μm for images B-F and H, and 2 μm x 2 μm for image G. Z-range 80 Å. When present, DYS R16-21 subphase concentration is 0.03 μM , CHOL 15 %. LE, liquid-expanded phase; LC, liquid-condensed phase.

Figure 5: Characterization of DYS R16-21 at an initial lipid surface pressure of 30 mN/m. (A) Variations of the surface pressure ($\Delta\pi$) and of the ellipsometric angle ($\delta\Delta$) induced by DYS R16-21 adsorption on phospholipid (black) and phospholipids/CHOL monolayers (gray). (B-H) Typical AFM topographic images of monolayers as indicated on the panels. Scan size 5 μm x 5 μm for images B-G and 2 μm x 2 μm for image H. Z-range 80 Å. When present,

DYS R16-21 subphase concentration is 0.03 μ M and CHOL, 15 %. LE, liquid-expanded phase; LC, liquid-condensed phase.

Figure 6: Molecular modeling of DYS R16-21. The 3D structure of DYS R16-21 was created in I-TASSER. The hydrophobic potential at the molecular surface (top row) is colored by increasing hydrophobicity from green (polar) to yellow (hydrophobic). The electrostatic potential at the molecular surface (middle row) is colored by increasing levels from red (-) to blue (+). The N-terminus (R16) is located at the left of the molecules. Left and right images in the top rows are representations of a 180°-rotation along the molecule's long axis. Bottom row: 3D structure of DYS R16-21 in cartoon representation. Each dystrophin repeat is a different grey. Amino acids of the CRAC (cholesterol recognition/interaction amino acid consensus) patterns are red.

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ACKNOWLEDGMENT

Supported by grants from AFM « association française contre les myopathies », GIS BRESMAT-Conseil Régional de Bretagne, and FEDER, « Fond Européen pour le Développement Régional ». SAL was funded by the « Ministère de l'Enseignement Supérieur et de la Recherche », AN was funded by the CNRS.

Authors thanks Biosit - Structure Fédérative de Recherche en Biologie-Santé de Rennes for circular dichroism facilities. Juliana Berland post-edited the English style.

ABBREVIATIONS

DOPC dioleoylphosphatidylcholine

DPPC dipalmitoylphosphatidylcholine

DOPS dioleoylphosphatidylserine

CHOL cholesterol

LE liquid expanded

LC liquid condensed

AFM atomic force microscopy

DYS dystrophin

β -DG β -dystroglycan

CRAC cholesterol recognition/interaction amino acid consensus

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